Outline	Computational biology vs bioinformatics	Bioinformatics databases	Hands-on
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Computational Molecular Biology and Bioinformatics

Introduction to Computational Biology and Bioinformatics

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Computational biology vs bioinformatics

2 Different areas of bioinformatics

- Sequence analysis
- Expression analysis
- Genetic analysis
- Epigenetic analysis
- System-level analysis
- Pathway analysis
- Bioinformatics databases





Computational biology vs bioinformatics

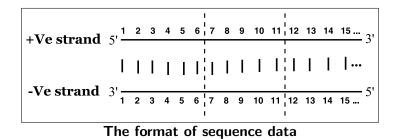
Computational biology (case \rightarrow model) is the study of biology systems using computational models techniques. The goal is to learn new biology, gain knowledge about living systems.

Bioinformatics (data \rightarrow information) is the creation of tools like statistical methods, algorithms, databases, etc. that solves problems. The goal is to build useful tools that successfully work on the biological data.

Note: Computational biology is 'science' whereas bioinformatics is 'engineering'.

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The sequence (or any subsequence) is always retrieved from the 5' to the 3' end, irrespective of the strand.

Note: 'A and T' and 'G and C' are complementary to each other in both these strands side by side.

Sequence annotation

Database 1: Genomic sequence

```
> ...
AATTCCGCGA ...
```

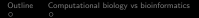
5'	1	2	3	4	5	6	7	8	9	10	 3'
5' +Ve	Α	A	Т	Т	C	C	G	C	G	A	
-Ve	Т	Т	Α	Α	G	G	С	Т	С	Т	
-Ve 3'	1	2	3	4	5	6	7	8	9	10	 5'

Database 2: Gene list

Gene Chrom Strand Start End Gene 1 chr1 + 2 6 Gene 2 chr1 - 5 8

Sequence of Gene1: ATTCC Sequence of Gene2: GCGG

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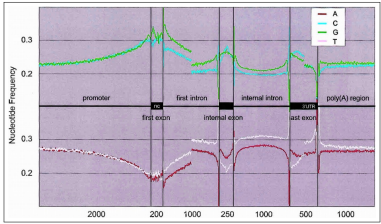
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Hands-on

Distribution of nucleotides in DNA



The frequencies of nucleotides throughout the genome

Various types of features are interesting in sequence analysis:

- *k*-mers: The frequency counts of subsequences of length *k*.
- Palindromes: The palindromic subsequences.
- CpG islands: The regions consisting of high density of CG and G+C contents, i.e. CpG observed by expected ratio and cumulative frequency of C and G, respectively.
- Special patterns: Special subsequence patterns.

<u>Note</u>: Several important genomic regions (e.g., promoter, gene body, etc.) have their unique features.

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FASTA and FASTQ formats

Notably, the alphabet size of DNA sequences (e.g., genes) is 4, whereas it is 20 for the amino acid sequences (e.g., proteins).

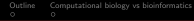
>gi|31563518|ref|NP_852610.1| microtubule-associated proteins 1A/1B light chain 3A isoform b [Homo sapiens] MKMRFFSSPCGKAAVDPADRCKEVQQIRDQHPSKIPVIIERYKGEKQLPVLDKTKFLVPDHVNMSELVKI IRRRLQLNPTQAFFLLVNQHSMVSVSTPIADIYEQEKDEDGFLYMVYASQETFGF

The sequence of a protein in FASTA format

```
@SEQ_ID
GATTTGGGGTTCAAAGCAGTATCGATCAAATAGTAAATCCATTTGTTCAACTCACAGTTT
+
!''*(((((***+))%%%++)(%%%%).1***-+*''))**55CCF>>>>>CCCCCCC65
```

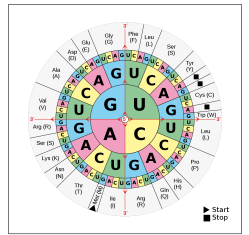
The sequence of a gene in FASTQ format

Note: In contrast to FASTA sequences, which includes only the identifier and sequence lines of size 50, a FASTQ sequence additionally takes the quality scores.



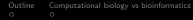
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The codon table



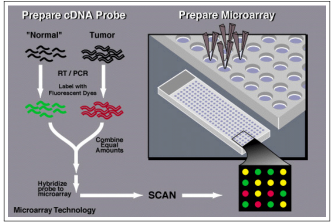
The conversion from RNA codons to proteins

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Microarray profiling



Profiling the cDNA microarrays

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Snapshot of expression data

Gene	ID	t1	t2	t3	t4	t5	t6	t7
G1		1.2	1.9	2.4	3.2	1.1	5.7	7.4
G2		3.2	3.9	4.4	5.3	3	7.8	9.5
G3		1	2.1	3.2	6.2	7.3	8.5	3.7
G1000		2.2	3.1	6.3	5.3	8.2	2.5	4.3

An example of expression dataset

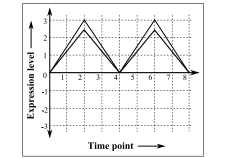
The rows and columns in an expression dataset can be different biomolecules (genes, proteins, microRNAs, etc.) and samples (time points, tissues, patients, etc.), respectively.

Different approaches of analysis – co-expression, differential expression, differential co-expression, co-expression dynamics, etc.



Definition (Co-expression)

Pairwise similarity pattern (spatial or temporal) of expression vectors.



Co-expression between a pair of expression vectors

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Differential expression

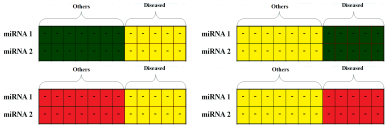
Gene	ID	c1	c2	c3	c4	n1	n2	n3	n4	n5	n6
G1		1.2	1.6	1.8	1.1	1	2	1.3	4	2	1.1
G2		1.1	1.5	1.3	1.8	2.1	1.1	1.1	1.1	2.3	1.5
G3		1.2	1.7	1.8	1.1	2	1.1	2.1	0.8	1.1	1.9
G100		2.1	1.6	1.7	1.4	2.3	2.7	2.8	2.9	1.3	2.1

Differential expression denotes varying patterns (spatial or temporal) of expression vectors between different phenotypes.

Differential co-expression

Definition (Differential co-expression)

Pairwise varying dependence (spatial or temporal) between expression vectors in different phenotypes.



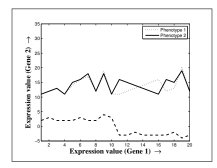
Different types of differential co-expression patterns

It can be mainly of two types – on/off case and gap/subsitution case.

Co-expression dynamics

Definition (Co-expression dynamics)

Pairwise varying dependence (spatial or temporal) between expression vectors based on another expression vector.



Genetic analysis is the overall process of studying and researching in fields of science that involve genetics and molecular biology.

Different areas of epigenetic analysis are listed below:

- Genomics
- Gene finding
- Phylogenetic analysis
- Genome wide association studies (GWAS)
- Proteomics
- Pharmacogenetics

Note: The environment has a major impact over the genome.

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The term epigenetics refers to heritable changes in gene expression (active versus inactive genes) that does not involve changes to the underlying DNA sequence; a change in phenotype without a change in genotype.

Different areas of epigenetic analysis are listed below:

- DNA methylation
- Histone modification

A methylated sequence (showing methylation with a '*') might appear as follows

ATCCCGACTGCAT

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System-level analysis

Molecular networks

- Protein-protein interaction networks
- Metabolic networks
- Regulatory networks TF-gene networks
- Post-translational networks Kinase-substrate networks
- RNA networks TF-miRNA Networks, miRNA-gene networks

Phenotypic networks

- Co-expression networks
- Genetic networks
- Disease networks



Biological pathway diagrams are used to describe molecular biology processes in a graphical way. A pathway is a set of related reactions in a given context, e.g., glycolysis, Krebs cycle, apoptosis, etc.

The role of bioinformatics on the pathway representations are creating specific requirements for their creation and curation.

Some popular pathway analysis tools are: KEGG, VisANT, etc.

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Bioinformatics databases

- Reference genome NCBI (http://www.ncbi.nlm.nih.gov)
- Genes and proteins NCBI, EMBL-EBI (https://www.ebi.ac.uk)
- MicroRNAs miRBase (http://www.mirbase.org)
- Other RNAs RNAdb (http://research.imb.uq.edu.au/RNAdb)
- Other biomolecules UCSC Genome Browser (http://genome.ucsc.edu)
- GO analysis Funcassociate 2.0 (http://llama.mshri.on.ca/funcassociate)
- Assembly conversion Galaxy (http://main.g2.bx.psu.edu)
- ID conversion DAVID (http://david.abcc.ncifcrf.gov/conversion.jsp)
- Gene enrichment analysis ShinyGo (http://bioinformatics.sdstate.edu/go)

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- Download the reference genome of SARS-CoV-2, the virus attributed to COVID-19, from NCBI following the steps below:
 - Open the NCBI portal.
 - Select "Assembly" and search for "SARS-CoV-2".
 - Download the complete reference genome sequence (RefSeq) in FASTA format.
 - **(**) Find out the frequencies of k-mers (k = 1, 2, 3) in the reference genome.
 - Identify the proteins that are already characterized in SARS-CoV-2.
 - Figure out from where you can download genome and protein sequences, annotation and a data report for all complete genomes of SARS-CoV-2.